Black Box Variational Inference with a Deterministic Objective

Faster, More Accurate, and Even More Black Box

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Problem statement

We all want to do accurate Bayesian inference quickly:

- In terms of compute (wall time, model evaluations, parallelism)
- In terms of analyst effort (tuning, algorithmic complexity)

Markov Chain Monte Carlo (MCMC) can be straightforward and accurate but slow.

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Black Box Variational Inference (BBVI) can be faster alternative to MCMC. But...

- ullet BBVI is cast as an optimization problem with an intractable objective \Rightarrow
- ullet Most BBVI methods use **stochastic gradient** (SG) optimization \Rightarrow
 - SG algorithms can be hard to tune
 - Assessing convergence and stochastic error can be difficult
 - SG optimization can perform worse than second-order methods on tractable objectives
- ullet Many BBVI methods employ a mean-field (MF) approximation \Rightarrow
 - Posterior variances are poorly estimated

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 - · Posterior variances are poorly estimated

Our proposal: replace the intractable BBVI objective with a fixed approximation.

- Better optimization methods can be used (e.g. true second-order methods)
- Convergence and approximation error can be assessed directly
- Can correct posterior covariances with linear response covariances
- This technique is well-studied (but there's still work to do in the context of BBVI)

⇒ Simpler, faster, and better BBVI posterior approximations ... in some cases.

Outline

- BBVI Background and our proposal
 - Automatic differentiation variational inference (ADVI) (a BBVI method)
 - Our approximation: "Deterministic ADVI" (DADVI)
 - Linear response (LR) covariances
 - Estimating approximation error
- Experimental results: DADVI vs ADVI
 - DADVI converges faster than ADVI, and requires no tuning
 - DADVI's posterior mean estimates' accuracy are comparable to ADVI
 - DADVI+LR provides more accurate posterior variance estimates than ADVI
 - DADVI provides accurate estimates of its own approximation error
 - But stochastic ADVI often results in better objective function values (eventually)
- · Sketch of theory and shortcomings

Notation

Data: y

Likelihood: $\mathcal{P}(y|\theta)$ Parameter: $\theta \in \mathbb{R}^{D_{\theta}}$

Prior: $\mathcal{P}(\theta)$ (density w.r.t. Lebesgue $\mathbb{R}^{D_{\theta}}$, nonzero everywhere)

We will be interested in means and covariances of the posterior $\mathcal{P}(\theta|y)$.

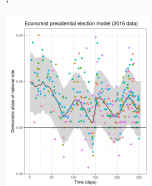
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Example: Election modeling (2016 US POTUS)

 $\mathsf{Data}\ y: \ \mathsf{Polling}\ \mathsf{data}\ (\mathsf{colored}\ \mathsf{dots})$

Likelihood $\mathcal{P}(y|\theta)$: Time series with random effects

Parameter θ : 15,098-dimensional

Interested in: Vote share on election day

MCMC time: 643 minutes (PyMC3 NUTS)

How can we approximate the posterior more quickly? One answer: variational inference.

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Variational inference [Blei et al., 2016]

We want the posterior $\mathcal{P}(\theta|y)$. Let $\mathrm{KL}\left(\mathcal{Q}(\theta)||\mathcal{P}(\theta)\right)$ denote KL divergence:

$$\mathrm{KL}\left(\mathcal{Q}(\theta)||\mathcal{P}(\theta)\right) = \underset{\mathcal{Q}(\theta)}{\mathbb{E}}\left[\log\mathcal{Q}(\theta)\right] - \underset{\mathcal{Q}(\theta)}{\mathbb{E}}\left[\log\mathcal{P}(\theta)\right].$$

The KL divergence is zero if and only if the two distributions are the same.

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A tautology:
$$\mathcal{P}(\theta|y) = \underset{\mathcal{Q}}{\operatorname{argmin}} \operatorname{KL} (\mathcal{Q}(\theta)||\mathcal{P}(\theta|y))$$

$$\text{Variational inference:} \qquad \mathring{\mathcal{Q}}(\theta) = \operatorname*{argmin}_{\mathcal{Q} \in \Omega_{\mathcal{Q}}} \operatorname{KL} \left(\mathcal{Q}(\theta) || \mathcal{P}(\theta|y) \right) \quad \dots \text{ for restricted } \Omega_{\mathcal{Q}}$$

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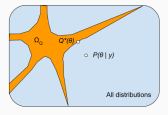
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We hope to choose $\Omega_{\mathcal{Q}}$ so that

- $\bullet \ \ \, \text{The optimization problem is tractable} \\ \to \ \, \text{simple} \, \Omega_{\mathcal{Q}} \, \, \text{are better}$
- The best approximation is a good one \rightarrow complex $\Omega_{\mathcal{Q}}$ are better

The approximation can be poor because

- Poor optimization
- \bullet The family $\Omega_{\mathcal{Q}}$ isn't expressive enough

Black-box variational inference

VI needs to solve:
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Involves the intractable integral $\underset{\mathcal{Q}(\theta)}{\mathbb{E}} [\log \mathcal{P}(\theta, y)]$

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How can we optimize this objective? Black-box VI [Ranganath et al., 2014]:

- Parameterize the family $\Omega_{\mathcal{Q}}$ using $\eta \in \mathbb{R}^{D_{\eta}}$ (so we have $\mathcal{Q}(\theta|\eta)$)
 - We will study **ADVI**, which takes $Q(\theta|\eta)$ to be Gaussian [Kucukelbir et al., 2017].
- Re-write the objective (using the reparameterization trick) as

$$\operatorname*{argmin}_{\eta} F(\eta) \quad \text{ where } \quad F(\eta) := \mathop{\mathbb{E}}_{\mathcal{N}_{\mathrm{std}}(z)} \left[f(\eta, z) \right].$$

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- ullet Use stochastic approximations to the gradient to minimize the exact $F(\eta)$
 - "Stochastic gradient" (used by almost all the BBVI community)
- Minimize a stochastic approximations to $F(\eta)$ (with exact gradients thereof)
 - The "sample average approximation" (our proposal)

Consider
$$\overset{*}{\mathcal{Q}}(\theta) = \operatorname*{argmin}_{\eta} F(\eta)$$
 where $F(\eta) := \underset{\mathcal{N}_{\mathrm{std}}(z)}{\mathbb{E}} [f(\eta, z)]$.

Let
$$\mathcal{Z}_N = \{z_1, \dots, z_N\} \stackrel{iid}{\sim} \mathcal{N}_{\mathrm{std}}(z)$$
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Algorithm 1

Stochastic gradient (SG)

ADVI (and most BBVI)

Fix
$$N$$
 (typically $N=1$)
 $t \leftarrow 0$
while Not converged do
 $t \leftarrow t+1$
Draw \mathcal{Z}_N
 $\Delta_S \leftarrow \nabla_\eta \ \hat{F}(\eta_{t-1}|\mathcal{Z}_N)$
 $\alpha_t \leftarrow \operatorname{SetStepSize}(\operatorname{Past\ state})$
 $\eta_t \leftarrow \eta_{t-1} - \alpha_t \Delta_S$
AssessConvergence(Past state)
end while
return η_t or $\frac{1}{M} \sum_{t'=t-M}^{t} \eta_{t'}$

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Algorithm 2

Sample average approximation (SAA)
Deterministic ADVI (DADVI) (proposal)

Fix N (our experiments use N=30) Draw \mathcal{Z}_N $t\leftarrow 0$ while Not converged do $t\leftarrow t+1$ $\Delta_D\leftarrow \mathrm{GetStep}(\hat{F}(\cdot|\mathcal{Z}_N),\eta_{t-1})$ $\eta_t\leftarrow \eta_{t-1}+\Delta_D$ AssessConvergence($\hat{F}(\cdot|\mathcal{Z}_N),\eta_t$)
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Our proposal: Apply Algorithm 2 with the ADVI objective.

Take better steps, easily assess convergence, with less tuning.

For each of a range of models (next slide), we compared:

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Stochastic ADVI methods:

- Mean field ADVI: We used the PyMC implementation of ADVI, together with its default termination criterion (based on parameter differences).
- Full-rank ADVI: We used the PyMC implementation of full-rank ADVI, together with the default termination criterion for ADVI described above.
- RAABBVI: To run RAABBVI, we used the public package viabel, provided by Welandawe et al. [2022].

We terminated unconverged stochastic ADVI after 100,000 iterations.

We evaluated each method on a range of models.

Model Name	$Dim\ D_\theta$	NUTS runtime	Description
ARM	Median 5	median 39 seconds	A range of linear models,
(53 models)	(max 176)	(max 16 minutes)	GLMs, and GLMMs
Microcredit	124	597 minutes	Hierarchical model with
			heavy tails and zero
			inflation
Occupancy	1,884	251 minutes	Binary regression with
			highly crossed random
			effects
Tennis	5,014	57 minutes	Binary regression with
			highly crossed random
			effects
POTUS	15,098	643 minutes	Autoregressive time series
			with random effects

Table 1: Model summaries.

Comparisons

To form a common scale for the accuracy of the posteriors, we report:

$$\varepsilon_{\text{METHOD}}^{\mu} := \frac{\mu_{\text{METHOD}} - \mu_{\text{NUTS}}}{\sigma_{\text{NUTS}}} \qquad \quad \varepsilon_{\text{METHOD}}^{\sigma} := \frac{\sigma_{\text{METHOD}} - \sigma_{\text{NUTS}}}{\sigma_{\text{NUTS}}}.$$

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We measure computational cost using both

- Wall time and
- Number of model evaluations (gradients, Hessian-vector products).

We compare achieved objective values using a large number of independent samples.

We report objective values and computation cost relative to DADVI.

Posterior mean accuracy

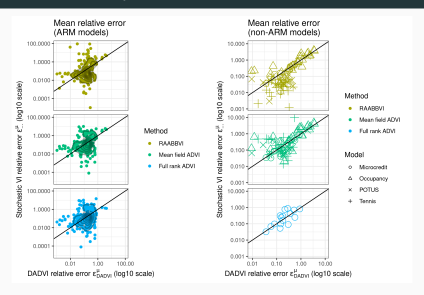
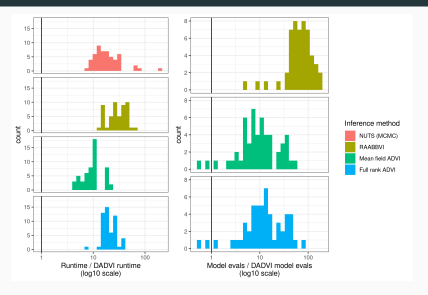


Figure 1: Posterior mean accuracy (relative to MCMC posterior standard deviation). Each point is a single named parameter in a single model. Points above the diagonal line indicate better DADVI or LRVB performance.

Computational cost for ARM models



 $\begin{tabular}{ll} Figure 2: Runtimes and model evaluation counts for the ARM models. Results are reported divided by the corresponding value for DADVI. \\ \end{tabular}$

Computational cost for non-ARM models

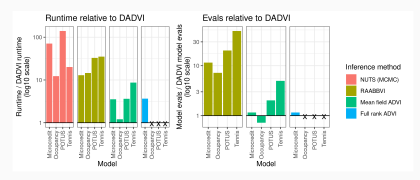


Figure 3: Runtimes and model evaluation counts for the non-ARM models. Results are reported divided by the corresponding value for DADVI. Missing model / method combinations are marked with an X.

Optimization traces for ARM models

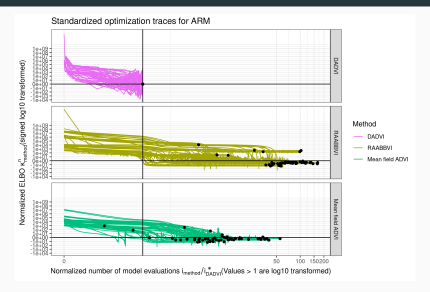


Figure 4: Optimization traces for the ARM models. Black dots show the termination point of each method. Dots above the horizontal black line mean that DADVI found a better ELBO. Dots to the right of the black line mean that DADVI terminated sooner in terms of model evaluations.

Optimization traces for non-ARM models

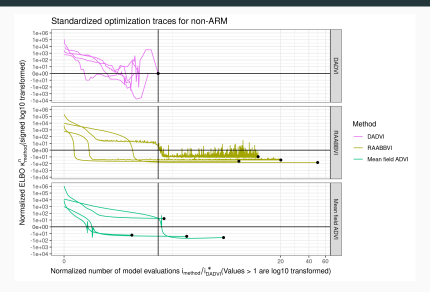


Figure 5: Traces for non-ARM models. Black dots show the termination point of each method. Dots above the horizontal black line mean that DADVI found a better ELBO. Dots to the right of the black line mean that DADVI terminated sooner in terms of model evaluations.

Experiment summary

⇒ DADVI is faster, simpler, and the posterior means are not worse.

But DADVI can additionally provide:

- Simple estimates of approximation error
- Improved (LR) posterior covariance esimates

Intractable objective:

$$\overset{*}{\eta} = \mathop{\mathrm{argmin}}_{\eta \in \mathbb{R}^{D_{\eta}}} \underset{\mathcal{N}_{\mathrm{std}}(z)}{\mathbb{E}} [f(\eta, z)]$$

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Answer: The same as a that of any M-estimator: asymptotically normal (as N grows)

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Linear response (LR) covariances improve covariance estimates by computing sensitivity of the variational means to particular perturbations. [Giordano et al., 2018]

Example: With a correlated Gaussian $\mathcal{P}(\theta|y)$, the ADVI means are exactly correct, the ADVI variances are underestimated, and LR covariances are exactly correct.

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Both DADVI error and LR covariances can be computed from the DADVI objective.

Stochastic ADVI does not produce an actual optimum of any tractable objective, so LR and M-estimator computations are unavailable.

Posterior standard deviation accuracy

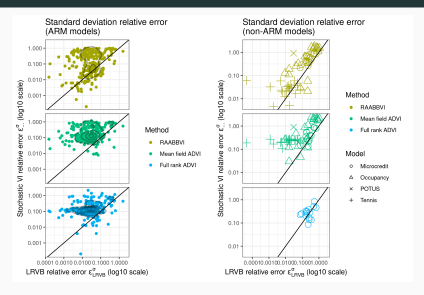


Figure 6: Posterior sd relative accuracy. Each point is a single named parameter in a single model. Points above the diagonal line indicate better DADVI or LRVB performance.

DADVI approximation error accuracy



Figure 7: Density estimates of $\Phi(\varepsilon^\xi)$ for difference models. All the ARM models are grouped together for ease of visualization. Each panel shows a binned estimate of the density of $\Phi(\varepsilon^\xi)$ for a particular model and number of draws N. Values close to one (a uniform density) indicate good frequentist performance. CG failed for the Occupancy and POTUS models with only 8 draws, possibly indicating poor optimization performance with so few samples.

Theory and shortcomings (sketch)

- SAA is hardly ever used for BBVI.¹ Why not?
 - Optimization literature predicts poor dimension dependence [Nemirovski et al., 2009]
 - Our experiments show that these results are too pessimistic
 - We prove better dimension dependence for SAA in some simple statistics models
 - A gap remains between theory and experiment.

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 - A gap remains between theory and experiment.
- Can I use SAA for very expressive approximating families?
 - We prove that the SAA fails for high-dimensional full-rank Gaussian approximations
 - We conjecture analogous problems with other expressive approximations

Given the benefits of DADVI over ADVI, there's more work to be done!

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Conclusion

Black Box Variational Inference with a Deterministic Objective: Faster, More Accurate, and Even More Black Box.

Giordano, R.*, Ingram, M.*, Broderick, T. (* joint first authors), 2023. (Arxiv preprint here.)

- By fixing the randomness in the ADVI objective, DADVI provides BBVI that is easier to use, faster, and more accurate than stochastic gradient.
- The approximation used by DADVI will not work in high dimensions for sufficiently expressive approximating distributions (e.g., full-rank ADVI).
- There appears to be a gap between the optimization literature and BBVI practice in high dimensions for a class of practically interesting problems.

References

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Supplemental material

Previous theoretical results

Intractable objective:

SAA approximation (DADVI):

$$\mathring{\eta} = \mathop{\rm argmin}_{\eta \in \mathbb{R}^{D_{\eta}}} \mathop{\mathbb{E}}_{\mathcal{N}_{\rm std}(z)} [f(\eta, z)]$$

$$\hat{\eta}(\mathcal{Z}_N) = \underset{\eta \in \mathbb{R}^{D_{\eta}}}{\operatorname{argmin}} \frac{1}{N} \sum_{n=1}^N f(\eta, z_n).$$

The idea of optimizing \hat{F} instead of SG on F is old and well-studied in the optimization literature, where \hat{F} is known as the **Sample average approximation (SAA)**.

Yet SAA is rarely used for BBVI.² One possible reason is the following:

Theorem [Nemirovski et al., 2009]: In general, the error of both SG and SAA scale as $\sqrt{D_{\theta}/N}$, where, for SG, N is the *total number of samples used*.

²Some exceptions I'm aware of: Giordano et al. [2018, 2022], Wycoff et al. [2022], Burroni et al. [2023].

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- For SG, each z_n gets used once (for a single gradient step)
- For SAA, each z_n gets used once per optimization step (of which the are many).
- Often, in higher dimensions, SAA requires more optimization steps.

Corollary: [Kim et al., 2015] In general, for a given accuracy, the computation required for SAA scales worse than SG as the dimension D_{θ} grows.

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But we got good results with D_{θ} as high as 15,098 using only only N=30. Why?

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Some first steps

Theorem [Giordano et al., 2023]: When $\mathcal{P}(\theta|y)$ is multivariate normal, and we use the mean-field Gaussian approximation, then, for any particular entry η_d of η , then $\left|\hat{\eta}_d - \mathring{\eta}_d\right| = O_p(N^{-1/2})$ irrespective of D_θ .

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Theorem [Giordano et al., 2023]: Assume $\mathcal{P}(\theta|y)$ has a "global-local" structure:

$$\theta = (\gamma, \lambda_1, \dots, \lambda_{D_{\lambda}}) \qquad \qquad \mathcal{P}(\gamma, \lambda_1, \dots, \lambda_{D_{\lambda}} | y) = \prod_{d=1}^{D_{\lambda}} \mathcal{P}(\gamma, \lambda_d | y).$$

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Under regularity conditions, the DADVI error scales as $\sqrt{\log D_{\lambda}/N}$, not $\sqrt{D_{\lambda}/N}$.

Proposal: The "in general" analysis of [Nemirovski et al., 2009] is too general for many practically interesting BBVI problems.

A negative result for expressive approximations

Theorem [Giordano et al., 2023]: Assume that $N < D_{\theta}$, and that we use a full-rank Gaussian approximation. Then the DADVI objective is unbounded below, and optimization of the DADVI objective will approach a degenerate point mass at $\operatorname{argmax}_{\theta} \log \mathcal{P}(\theta|y)$.

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Proof sketch: For any value of the variational mean, the DADVI objective only depends on $\mathcal{P}(\theta|y)$ evaluated in a subspace spanned by \mathcal{Z}_N . The variational objective can be driven to $-\infty$ by driving the variance to zero in the subspace orthogonal to \mathcal{Z}_N .

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Proposal: All sufficiently expressive variational approximations (e.g. normalizing flows) will fail in the same way in high dimensions. However, this pathology can be obscured and overlooked in practice by low-quality optimization.

Linear response covariances

Posterior variances are often badly estimated by mean-field (MF) approximations.

Example: With a correlated Gaussian $\mathcal{P}(\theta|y)$ with ADVI, the ADVI means are correct, but the ADVI variances are underestimated.

Take a variational approximation $\mathring{\eta}:= \operatorname{argmin}_{\eta \in \mathbb{R}^{D_{\eta}}} \operatorname{KL}_{\operatorname{VI}}(\eta)$. Often,

$$\underset{\mathcal{Q}(\theta|_{\eta}^{*})}{\mathbb{E}}[\theta] \approx \underset{\mathcal{P}(\theta|y)}{\mathbb{E}}[\theta] \quad \text{but} \quad \underset{\mathcal{Q}(\theta|_{\eta}^{*})}{\text{Var}}(\theta) \neq \underset{\mathcal{P}(\theta|y)}{\text{Var}}(\theta). \tag{1}$$

Example: Correlated Gaussian $\mathcal{P}(\theta|y)$ with ADVI.

Linear response covariances use the fact that, if $\mathcal{P}(\theta|y,t) \propto \mathcal{P}(\theta|y) \exp(t\theta)$, then

$$\frac{d \underset{\mathcal{P}(\theta|y,t)}{\mathbb{E}} [\theta]}{dt} = \underset{t=0}{\text{Cov}} (\theta).$$
 (2)

Let $\mathring{\eta}(t)$ be the variational approximation to $\mathcal{P}(\theta|y,t)$, and take

$$\operatorname{LRCov}_{\mathcal{Q}(\theta|\mathring{\eta})}(\theta) = \frac{d \mathbb{E}_{\mathcal{Q}(\theta|\mathring{\eta}(t))}[\theta]}{dt} = \left(\nabla_{\eta} \mathbb{E}_{\mathcal{Q}(\theta|\mathring{\eta})}[\theta]\right) \left(\nabla_{\eta}^{2} \operatorname{KL}_{\operatorname{VI}}(\mathring{\eta})\right)^{-1} \left(\nabla_{\eta} \mathbb{E}_{\mathcal{Q}(\theta|\mathring{\eta})}[\theta]\right)$$

Example: For ADVI with a correlated Gaussian $\mathcal{P}(\theta|y)$, $\operatorname{LRCov}_{\mathcal{Q}(\theta|\eta)}(\theta) = \operatorname{Cov}_{\mathcal{Q}(\theta|\eta)}(\theta)$.